

APPENDIX C. FORM FDA 1572

						Form American CAIF 11- 2012 201	
DEP	ARTMENT OF H FOOD AND	DRUG ADMINI		VICES	- 1	Form Approved: OMB No. 0910-0014. Expiration Date: January 31, 2006. See OMB Statement on Reverse.	
(ΤΙ TLE 21,	STATEMEN CODE OF FEDE (See instru		ATIONS (CFR			NOTE: No investigator may participate in al investigation until he/she provides the spons a completed, signed Statement of investigat Form FDA 1572 (21 CFR 312.53(c)).	or wt
1. NAME AND ADDRES	S OF INVESTIGATO	R			<u>'</u>		
2. EDUCATION, TRAINI	NG, AND EXPERIEN	NCE THAT QUAL	IFIES THE INVES	STIGATOR AS A	AN EXPERT	IN THE CLINICAL INVESTIGATION OF THE	
DRUG FOR THE USE	_			_			
	□cu	IRRICULUM VITA	AE	OTHER	RSTATEMEN	NT OF QUALIFICATIONS	
NAME AND ADDRES	S OF ANY MEDICAL	L SCHOOL, HOS	PITAL OR OTHE	R RESEARCH F	ACILITY WE	HERE THE CLINICAL INVESTIGATION(S) WILL	
BE CONDUCTED.		·					
NAME AND ADDRES	S OF ANY CLINICAL	LABORATORY	FACILITIES TO E	BE USED IN TH	E STUDY.		
5. NAME AND ADDRES	S OF THE INSTITUT	TIONAL REVIEW	BOARD (IRB) TH	HAT IS RESPON	SIBLE FOR	REVIEW AND APPROVAL OF THE STUDY(IES).
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B. ATTACH THE FOLLOWING CLINICAL PRO	OTOCOL INFORMATION:
	A GENERAL OUTLINE OF THE PLANNED INVESTIGATION INCLUDING THE ESTIMATED DURATION OF JUMBER OF SUBJECTS THAT WILL BE INVOLVED.
SUBJECTS TO BE TREATED WITH INVESTIGATED; CHARACTERISTIC	ONS, AN OUTLINE OF THE STUDY PROTOCOL INCLUDING AN APPROXIMATION OF THE NUMBER OF ITHE DRUG AND THE NUMBER TO BE EMPLOYED AS CONTROLS, IF ANY; THE CLINICAL USES TO BE OS OF SUBJECTS BY AGE, SEX, AND CONDITION; THE KIND OF CLINICAL OBSERVATIONS AND DUCTED; THE ESTIMATED DURATION OF THE STUDY; AND COPIES OR A DESCRIPTION OF CASE
9. COMMITMENTS:	
	coordance with the relevant, current protocol(s) and will only make changes in a protocol after notitying the rotect the safety, rights, or welfare of subjects.
I agree to personally conduct or super	rvise the described investigation(s).
	persons used as controls, that the drugs are being used for investigational purposes and I will ensure that Informed consent in 21 CFR Part 50 and institutional review board (IRB) review and approval in 21 CFI
I agree to report to the sponsor advers	se experiences that occur in the course of the investigation(s) in accordance with 21 CFR 312.64.
I have read and understand the inform	nation in the investigator's brochure, including the potential risks and side effects of the drug.
I agree to ensure that all associates, in meeting the above commitments.	colleagues, and employees assisting in the conduct of the study(les) are informed about their obligation
I agree to maintain adequate and accardance with 21 CFR 312.68.	curate records in accordance with 21 CFR 312.62 and to make those records available for inspection
	. I also agree to promptly report to the IRB all changes in the research activity and all unanticipate ubjects or others. Additionally, I will not make any changes in the research without IRB approval, exce
problems involving risks to human su where necessary to eliminate apparen	ubjects or others. Additionally, I will not make any changes in the research without IRB approval, except nt immediate hazards to human subjects. Trements regarding the obligations of clinical investigators and all other pertinent requirements in 21 CF
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